Glucose-induced Downregulation of Angiotensin II and Arginine Vasopressin Receptors in Cultured Rat Aortic Vascular Smooth Muscle Cells
Role of Protein Kinase C

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Abstract

Early diabetes mellitus is characterized by impaired responses to pressor hormones and pressor receptor downregulation. The present study examined the effect of elevated extracellular glucose concentrations on angiotensin II (AII) and arginine vasopressin (AVP) receptor kinetics in cultured rat vascular smooth muscle cells (VSMC). Scatchard analysis of [3H]AVP and 125I-AII binding to confluent VSMC showed that high glucose concentrations (20 mM) similarly depressed AVP and AII surface receptor Bmax but did not influence receptor Kd. This receptor downregulation was not reproduced by osmotic control media containing either L-glucose or mannitol. Receptor downregulation was maximal at a glucose concentration of 15–20 mM and required 24–48 h for a maximum effect. Normalization of the extracellular glucose concentration allowed complete recovery of AVP and AII binding within 48 h. Receptor downregulation was associated with depressed AVP and AII-stimulated intracellular signaling and cell contraction. High glucose concentrations induced a sustained activation of protein kinase C (PKC) in VSMC, which was prevented by coincubation with H-7. H-7 also markedly attenuated glucose-induced downregulation of AVP and AII receptors on VSMC. This study demonstrates a novel cellular mechanism whereby high extracellular glucose concentrations directly and independently downregulate pressor hormone receptors and their function on vascular tissue via glucose-stimulated PKC activation. (J. Clin. Invest. 1992. 90:1992–1999.) Key words: diabetes mellitus • protein kinase C • vascular smooth muscle cells • microangiopathy • vascular injury

Introduction

Early diabetes mellitus in both experimental animals and man is associated with increased blood flow to many tissues, including the kidney, myocardium, retina, skin, muscle, and brain (1–7). These early changes in regional hemodynamics have in turn been strongly implicated in the pathogenesis of the widespread microvascular injury that characterizes this disease (8–10). The principal determinants of this deleterious increase in tissue perfusion is a reduction in arteriolar tone (11). Pressor hormones such as angiotensin II (AII)1 and arginine vasopressin (AVP) are important modulators of vascular tone (12) and studies of the diabetic rat have demonstrated an early reduction in the aortic contractile response to AII and epinephrine (13). Moreover, glomerular hemodynamic responses to AII are also markedly blunted in diabetic rats (14).

The aforementioned impaired response to pressor hormones could result from either impaired postreceptor actions of the hormone or reduced hormone-receptor binding. In keeping with the latter, reduced glomerular AII binding has been demonstrated in the diabetic rat (15, 16); reduced AVP binding to platelet V1 receptors has also been observed in patients with diabetes mellitus (17). Additional studies reveal a reduction in the number of beta-adrenergic receptors in experimental diabetes and platelet thromboxane A2 receptors in human diabetes (18, 19), suggesting that the diabetic state may be associated with a generalized reduction in the expression of a variety of pressor receptors.

In spite of the potential pathophysiological significance of pressor receptor downregulation with regard to the initiation and/or propagation of vascular injury in diabetes, the mechanisms responsible for these marked changes in surface receptor density are unknown. Recent evidence suggests that protein kinase C (PKC), a multifunctional Ca2+- and phospholipid-dependent serine/threonine kinase system, may play an important role in regulating the cell surface density of many receptors (20–22). In this regard, we have shown that hormone-activated PKC is an important component of homologous hormone receptor desensitization in cultured rat vascular smooth muscle cells (VSMC) (23, 24). Others report a similar PKC-dependent downregulation of a variety of receptors in different tissues (25–27). More recent reports suggest that PKC may modulate surface receptor density by regulating receptor mRNA expression and receptor biosynthesis (28). In view of the potentially important role of PKC in the regulation of receptor biosynthesis and expression, it is intriguing that high extracellular glucose concentrations have recently been shown to promote the activation of PKC in numerous tissues, including vascular tissue, both in vivo and in vitro (29–32). Together, these observations raise the possibility that glucose-induced PKC activation could provide a novel biochemical mechanism to account for downregulation of pressor receptors in patients with diabetes mellitus.

The present in vitro study thus examines the hypothesis that elevated extracellular glucose concentrations can directly promote the downregulation of AII and AVP receptors on VSMC and thereby impair the subsequent pressor response of VSMC to these hormones via mechanisms dependent on glucose-induced PKC activation in VSMC.

1. Abbreviations used in this paper: AII, angiotensin II; AVP, arginine vasopressin; DAG, diacylglycerol; PKC, protein kinase C; PSS, physiological saline solution; VSMC, vascular smooth muscle cells.
Methods

Materials. AVP, AII, and H7 were purchased from Sigma Chemical Co. (St. Louis, MO). 125I-AII and [3H]AVP were obtained from New England Nuclear (Wilmington, MA). 45Ca2+ was obtained from Amersham Corp. (Arlington Heights, IL). The VRKRTLRL peptide substrate was a generous gift from Drs. Lynn E. Heasley and Gary L. Johnson, National Jewish Center for Immunology and Respiratory Medicine, Denver, CO.

Preparation of cultured rat VSMC. Rat aortic VSMC were isolated and cultured using a modification of the method described by Chamley et al., as previously described from this laboratory (23, 24, 33). Briefly, under sterile conditions, aortas were resected from Sprague-Dawley rats and cleaned of adventitia and connective tissue. The vessels were chopped and incubated for 2 h in Eagle's MEM (Gibco Laboratories, Grand Island, NY) containing 2 mg/ml collagenase. The resulting cell suspension was plated on 35-mm dishes and grown in MEM containing 2 mM L-glutamine, 2 g/liter NaHCO3, 100 IU/ml penicillin, 100 mg/liter streptomycin, and 10% FCS at 37°C in 95% humidified air and 5% CO2. At confluence, VSMC from second through sixth passage were used.

Test media. To examine specifically the effects of various extracellular D-glucose concentrations on AVP or AII binding to VSMC and the subsequent biological and physiological response to these agents, VSMC were exposed to one of four test media: a) control medium: consisted of MEM supplemented with 10% FCS and containing 5 mM D-glucose; b) high glucose medium: identical to the control medium except that it was supplemented with D-glucose to increase the glucose concentration up to 30 mM; c) l-glucose osmotic control medium: identical to the control medium but supplemented with a cell permeable but poorly metabolized glucose isomer, l-glucose (25 mM); or d) mannitol osmotic control medium: identical to the control medium but supplemented with poorly diffusible hexose, mannitol (25 mM).

Receptor binding studies. The experiments examining AVP and AII binding to VSMC were performed using the radioligands 3H]AVP and 125I-AII as previously described (34). To determine AVP binding after preexposure to the test media for various time periods, confluent VSMC monolayers on 35 × 10-mm dishes were washed twice with ice-cold binding buffer (119.2 mM NaCl, 3 mM KCl, 1.2 mM MgSO4, 1 mM CaCl2, 1.2 mM KH2PO4, 10 mM glucose, 10 mM Hepes, 0.1% BSA, pH 7.4). The cells were then incubated with the same buffer containing 2 × 10−9 M [3H]AVP (specific activity 67.7 μCi/μmol) with or without unlabeled AVP for 90 min at 4°C to prevent receptor internalization and achieve saturation binding conditions. Binding was terminated and the unbound radioligand removed by rapid washing four times with 2 ml ice-cold binding buffer. The cells were then solubilized in 0.1% SDS and 0.1 N NaOH and the cell-associated radioactivity was determined by scintillation counting (Tri-carb 460C; Packard Instrument Co. Inc., Downers Grove, IL). Specific binding of AVP was defined as total binding (2 × 10−12 M [3H]AVP) minus nonspecific binding (2 × 10−9 M [3H]AVP and 10−8 M unlabeled AVP). A 50-μl aliquot of the solubilized cells was assayed for protein content by the method of Lowry. All binding studies were performed using similar conditions except that the composition of the binding buffer was 50 mM Tris-HCl, 100 mM NaCl (pH 7.4), 5 mM MgCl2, supplemented with 0.5 mg/ml bacitracin and 0.2% BSA. Specific 125I-AII binding to VSMC was defined as total binding (125I-AII, 50–100 fmol, specific activity 103 dpm/ml) minus nonspecific binding (unlabeled AII [10−8 M]). Each binding assay was performed in triplicate and binding constants were determined by Scatchard analysis of binding data fitted to a line.

Measurement of PKC activity in VSMC. For the measurement of in situ PKC activity in VSMC, a modification of the method recently described by Heasley and Johnson (35, 36) was used. VSMC were seeded into flat-bottomed 96-well microtiter plates at a density of 20,000/well. VSMC were maintained in regular growth medium for 1 to 3 d until confluent. The monolayer was then washed with 200 μl of Hanks' balanced salt solution before incubation with one of the four test media described above. After the desired exposure time, the test medium was aspirated and replaced with 40 μl of a buffered salt solution containing 137 mM NaCl, 5.4 mM KCl, 10 mM MgCl2, 0.3 mM sodium phosphate, 0.4 mM potassium phosphate, 25 mM β-glycerophosphate, 5.5 mM D-glucose, 5 mM EGTA, 1 mM CaCl2 (100 mM Ca2+-free), 100 μM [γ32P]ATP (~5,000 cpm/μmol), 50 μg/ml digitonin, and 20 mM Hepes (pH 7.2, 30°C). In addition, 100 μM of a PKC-specific peptide substrate (VRKRTLRL) was added to the buffer. This short synthetic peptide is based on the sequence surrounding a major PKC-dependent phosphorylation site within the epidermal growth factor (EGF) receptor (37). This peptide substrate is not phosphorylated by cyclic nucleotide-dependent or Ca2+/calmodulin-dependent protein kinases or 56 kinase and has been extensively characterized to be a highly specific substrate for PKC (35, 36). By permeabilizing VSMC with the digitonin contained in the buffer, the VRKRTLRL peptide enters VSMC along with [γ32P]ATP to allow a highly selective and rapid analysis of in situ PKC activity. The concentration of digitonin (50 μg/ml) used in this assay does not modify VSMC morphology or promote cell detachment. Moreover, PKC activity is retained by the monolayer postpermeabilization (PKC activity in the supernatant was <10% total PKC activity). This latter finding is consistent with the concept that the PKC activity being measured using this assay is tightly associated with the cell membrane as has been proposed for the active form of PKC (20, 38, 39). The kinase reaction was linear with time for up to 20 min and was therefore allowed to proceed for 10 min at 30°C before termination by the addition of 25% (wt/vol) TCA (final TCA concentration 5%)). Aliquots (45 μl) of the acidified reaction mixture were spotted onto 2-cm phosphocellulose circles (P81; Whatman Inc., Clifton, NJ) and washed batchwise three times with 75 mM phosphoric acid and one wash with 75 mM sodium phosphate (pH 7.5) (500 ml/2-min wash). Because of the basicity of the VRKRTLRL substrate, it is retained by the phosphocellulose filter at neutral pH, whereas contaminating [32P]ATP is removed. The PKC-dependent phosphorylation of the peptide substrate bound to the filter was quantified by scintillation counting. Results are expressed as PKC-dependent peptide phosphorylation, pmol/min per mg VSMC protein.

Measurement of 45Ca2+ efflux from VSMC. 45Ca2+ efflux studies were performed as previously described (24, 40, 41). Briefly, after exposure to the test media, confluent VSMC on 35-mm dishes were washed with physiological saline solution (PSS) and incubated with 1 ml fresh test medium containing 8 μCi 45Ca2+ (specific activity) for 3 h at 37°C to allow preloading VSMC with 45Ca2+. Thereafter, VSMC were rinsed rapidly (10 × 1 ml PSS in 60 s) and then incubated with 1 ml PSS. The PSS was removed and replaced with fresh PSS at 1-min intervals for 6 min. Samples at each time interval were placed into vials for liquid scintillation counting. After 6 min, the PSS was supplemented with either AVP (10−7 M) or AII (10−7 M) and VSMC were exposed to these agents for 1 min. 45Ca2+ released by VSMC during the incubation with these agents and for subsequent 5 × 1 min incubations with 1 ml PSS was quantified by liquid scintillation counting. The VSMC were solubilized as described above and the cell-associated radioactivity was counted. Total 45Ca2+ released after the addition of the effector minus basal release was expressed as a percentage of total available cellular radioactivity at the time of stimulation with the effector.

VSMC shape change studies. The functional response of VSMC to AVP or AII was assessed by quantitation of cell surface area changes using phase-contrast microscopy (IM35; Carl Zeiss, Inc., Oberkothen, Germany) and a computerized digital image analyzer (Zidas, Carl Zeiss Inc.) that had been calibrated using a micrometer scale, as previously reported from this laboratory (23, 24). For these studies, only sparsely plated primary cultures of VSMC in 35 × 10-mm dishes were used because VSMC may lose their contractility after subculture (42). After preexposure to the test media, VSMC were incubated for 20 min in the presence of fresh test media at 37°C. After this equilibration period, the area of individual cells was measured. A 20× objective was used for all studies on groups of 10–15 cells. The microscopic field was chosen randomly and not selected for cell shape or size. AVP (10−7 M)

Glucose-induced Receptor Downregulation 1993
or AII (10⁻⁴ M) was then added to the medium and the VSMC incubated for a further 15 min at 37°C. Changes in VSMC surface area in the same group of cells in response to AVP or AII was then measured with the image analyzer. Several precautions were incorporated into the protocol. Three measurements of each cell were taken and any group of measurements showing a standard deviation of > 20% of the mean was discarded. Spontaneous cell shape changes due to manipulation of the culture dishes in the absence of effector was measured at < 5% and thus only a value of ≥ 15% change in surface area was considered to be a significant response to AVP or AII. The method has been further validated in this laboratory by comparing the digital image-analyzed assessment of cell surface area changes with the measured area on photographs of the same cells. The results show excellent correspondence between the two methods.

Statistical analysis: Results are expressed as mean±SEM. Statistical analysis was made using the paired or unpaired Student’s t test or analysis of variance with a Bonferroni correction as appropriate. For each experiment, n refers to the number of studies (each in triplicate).

Results

Effect of high extracellular glucose concentrations on AVP and AII binding to VSMC. Studies examining AVP and AII binding to VSMC were performed under conditions at which saturation binding occurs for both ligands and at 4°C to prevent receptor internalization so that only surface binding was measured. Equilibration binding for both radioligands occurred by 60 min, remained stable over the next 30 min, and was identical in the presence of all test media. The binding characteristics of both radioligands were specific, time and protein dependent, saturable, and stable in VSMC up to sixth passage. For both AVP and AII, the nonspecific binding was always < 15% and was not affected by exposure to the different test media.

In the presence of the control medium (D-glucose 5 mM), Scatchard transformation of the binding data for specific [³H]AVP binding to confluent VSMC yielded a linear plot, demonstrating a single class of AVP-binding sites with a maximum number of binding sites (B_max) of 1.99 x 10⁻¹² mol/mg cell protein and a K_d of 2.15 x 10⁻⁹ mol (Fig. 1). In contrast, however, preexposing VSMC for 48 h to a high glucose medium (D-glucose 20 mM) markedly reduced AVP binding by almost 40% versus control medium, due to a significant decline in B_max (1.22 x 10⁻¹² mol/mg cell protein) with no significant change in K_d (2.26 x 10⁻⁹ mol) (Fig. 1).

Similarly, in the presence of control medium, AII binds to a single class of receptors on VSMC with a B_max of 2.96 x 10⁻¹³ mol/mg cell protein and a K_d of 3.2 x 10⁻⁹ mol (Fig. 1). However, preexposure of VSMC to a high glucose medium (d-glucose 20 mM) for 48 h resulted in a marked reduction in AII binding to VSMC. This was similar in magnitude to the effect of the high glucose environment on AVP binding to VSMC and was also due to a significant reduction in AII receptor B_max (2.27 x 10⁻¹³ mol/mg cell protein) with no significant change in K_d (3.36 x 10⁻⁹ mol). Thus exposing VSMC to a high extracellular glucose environment resulted in a significant decrease in AVP and AII binding to VSMC in both cases due to a similar decline in surface density of these pressor receptors with no significant change in receptor affinity. These changes did not reflect changes in VSMC protein content or cell number as both of these parameters were equivalent after a 48-h exposure of confluent VSMC to all test media. Furthermore, the glucose-induced decrease in hormone binding did not represent VSMC toxicity or accelerated cell death because cell detachment rates, percent lactate dehydrogenase release and percent trypan blue exclusion, were also similar after a 48-h exposure to all test media (data not shown).

To examine the possibility that changes in AVP or AII surface binding were related to the increased extracellular osmolality of the high glucose medium, the effects of a 48-h exposure to the two osmotic control media (L-glucose or mannitol) on AVP and AII binding to VSMC was examined. The results indicate that the downregulation of AVP and AII receptors on VSMC was specific for D-glucose and is not reproduced by either of the two osmotic test media (Table 1). Moreover, the fact that high concentrations of the poorly metabolized L-glucose isomer did not modify AVP or AII binding to VSMC suggests that intracellular metabolism of D-glucose is necessary for its effects on pressor receptor expression to develop.

Glucose concentration dependency of AVP and AII receptor downregulation in VSMC. The next study examined the relationship between extracellular glucose concentration and depression of AVP and AII binding to VSMC. Confluent VSMC were incubated with different D-glucose concentrations for 48 h before determining the binding of [³H]AVP or [¹²⁵]A-II to VSMC. The glucose concentration dependency of inhibition of AVP and AII binding to VSMC was a threshold effect with the specific binding of both hormones being significantly depressed at a glucose concentration of 15 mM and maximally depressed between 15 and 20 mM (Fig. 2).

Table I. Effect of a 40-h Incubation with the Different Test Media on the Specific Binding of [³H]AVP and [¹²⁵]A-II to Confluent VSMC

<table>
<thead>
<tr>
<th>Test media</th>
<th>AVP</th>
<th>All</th>
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<tbody>
<tr>
<td></td>
<td>% specific binding</td>
<td>% specific binding</td>
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<tr>
<td>Glucose (5 mM)</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>L-glucose control</td>
<td>103.9±5.4</td>
<td>101.0±6.1</td>
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<tr>
<td>Mannitol control</td>
<td>98.7±5.2</td>
<td>100.8±7.3</td>
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<tr>
<td>Glucose (20 mM)</td>
<td>61.8±4.9*</td>
<td>66.3±5.5*</td>
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Specific binding in the presence of control medium (glucose 5 mM) is designated 100% and results in the presence of the other test media are expressed as percentage of this control. * P < .01 vs control.
Figure 2. The glucose concentration dependency of AVP- and All-specific surface binding to confluent VSMC. VSMC were exposed to either control medium (glucose 5 mM) or identical medium supplemented with various concentrations of d-glucose for 48 h before determining the specific surface binding of AVP (open bars) and All (hatched bars). AVP or All binding in presence of control medium is designated 100% and binding in the presence of the other test media is expressed as percent of control. *P < 0.01 vs. control binding (n = 3).

Time dependency of glucose-induced downregulation of AVP and All receptors on VSMC. To examine the time dependency of the downregulation of AVP and All receptors in the presence of a high glucose concentration, VSMC were exposed to a glucose concentration of 20 mM for increasing time periods up to 48 h. Fig. 3 shows that > 12-h exposure to a high glucose concentration was required to significantly downregulate either the AVP or All receptor and between 24 and 48 h was required for maximal glucose-induced downregulation of these receptors on VSMC. The maximum time exposure to the high glucose medium in all studies was 48 h.

Recovery of AVP and All receptors. Having demonstrated the slow onset of glucose-induced pressor receptor downregulation, the next study examined receptor recovery after restoring the extracellular glucose concentration to normal. VSMC were exposed to a high d-glucose concentration (20 mM) for 48 h to induce maximal downregulation of AVP and All receptors. The high glucose medium was then replaced with control medium (glucose 5 mM) for various time periods before measuring specific AVP or All binding to VSMC. The characteristics of AVP and All receptor recovery were very similar. Both receptors showed signs of recovery within 12 h of normalizing the extracellular glucose concentration but required up to 48 h for full receptor recovery to occur (Fig. 4). These results demonstrate that glucose-induced effects on both AVP and All surface receptors on VSMC are similarly slow to develop and slow to recover. Furthermore, the complete reversibility of this phenomenon confirms that glucose-induced receptor downregulation does not involve any artifact due to cell death.

Glucose-induced activation of PKC in VSMC. The next series of studies were designed to explore whether PKC activation is involved in the mechanism(s) whereby glucose promoted the downregulation of pressor receptors on VSMC. PKC activation has been implicated in the regulation of hormone receptor biosynthesis and expression. Elevated extracellular glucose concentrations have been shown to induce the activation of PKC in cultured retinal endothelial cells and isolated glomeruli via an increased flux of glucose through an intracellular pathway culminating in the enhanced de novo synthesis diacylglycerol (DAG) (29, 32), which in turn promotes the activation of PKC (20, 38). The next series of studies thus examined whether elevated glucose concentrations activate PKC in VSMC. Fig. 5 shows that after a 3-h exposure to a high glucose (20 mM) medium, in situ PKC activity was markedly increased compared with the PKC activity measured in VSMC exposed to control medium (glucose, 5 mM). Of interest, the glucose-induced increase in PKC activity was sustained for up to 48 h, provided that the extracellular glucose concentration remained elevated. In contrast, 48-h exposure to either of the two osmotic control media produced no appreciable activation of PKC, confirming that PKC activation was not osmotically mediated but required the metabolism of glucose (data not shown). Further studies confirmed that PKC activation was glucose-concentration dependent. A threshold glucose concentration of 15 to 20 mM was required to induce a maximal PKC response to glucose (data not shown).

![Graph](image-url)
Phosphorylation was determined in confluent VSMC by measuring the phosphorylation of the PKC-specific VRKRTLRL peptide substrate. The results are expressed as pmol phosphate transferred to this substrate/min per mg VSMC protein. *P < 0.01 vs. the time-matched control (n = 4).

To examine the role of glucose-induced PKC activation in mediating pressor receptor downregulation, a means of manipulating PKC activation by glucose was necessary. The next study determined the capacity of H7, a relatively specific inhibitor of PKC activity (43), to prevent glucose-induced PKC activation. VSMC were coincubated with H7 (5 × 10⁻³ M in 0.1% DMSO) in either control medium or high glucose (20 mM) medium for 48 h before determining in situ PKC activity. DMSO (0.1%) alone had no effect on basal or glucose stimulated PKC activity (data not shown). With control medium, H7 reduced basal PKC activity. Moreover, when high glucose medium was coincubated with H7, glucose-induced PKC activation was almost completely prevented (Fig. 6). The fact that the phosphorylation of the VRKRTLRLR peptide substrate was markedly attenuated by a recognized PKC inhibitor (H7) demonstrates that the phosphorylating activity being measured in the permeabilized VSMC was specific for PKC.

The role of PKC in glucose-induced AVP and AI receptor downregulation in VSMC. After demonstrating that the sustained activation of PKC induced by chronic high glucose exposure could be prevented by coincubation with H7, the same experimental maneuver was used to examine specifically the role of PKC in mediating glucose-induced pressor receptor downregulation in VSMC. Fig. 7 shows that, in the absence of H7, 48-h exposure to high glucose (20 mM) medium resulted in a significant downregulation of AVP and AI receptors on VSMC. In contrast, when the high glucose medium was supplemented with H7 to prevent glucose-induced PKC activation, the downregulation of AVP and AI receptors was almost completely prevented. These results illustrate that glucose-induced downregulation of AVP and AI receptors is strongly dependent on the capacity of glucose to activate PKC in VSMC. Moreover, it is of interest that inhibition of basal PKC activity, in the presence of control medium, increased the expression of both AVP and AI receptors, suggesting that PKC activity is involved in regulating the basal expression of these receptors.

**Figure 5.** Glucose-induced PKC activity in VSMC. Confluent VSMC were exposed for varying time periods to control medium (glucose 5 mM) (open bars) or high glucose medium (glucose 20 mM) (spotted bars) before measuring PKC activity. PKC activity was measured in situ in digitonin-permeabilized VSMC by determining the phosphorylation of the PKC-specific VRKRTLRL peptide substrate. Results are expressed as pmol phosphate transferred to this substrate/min per mg VSMC protein. *P < 0.01 vs. the time-matched control (n = 4).

**Figure 6.** Inhibition of glucose-induced PKC activity by H7. Confluent VSMC were exposed to either control medium (glucose 5 mM) (open bars) or high glucose medium (glucose 20 mM) (spotted bars) + H7 (5 × 10⁻³ M) for 48 h before measuring in situ PKC activity (n = 3).

**Figure 7.** Effect of PKC inhibition on glucose-induced AVP and AI receptor downregulation in VSMC. Confluent VSMC were exposed to either control medium (glucose 5 mM) or high glucose medium (glucose 20 mM) ± H7 (5 × 10⁻³ M) for 48 h before determining the surface binding of AVP and AI to VSMC. Specific AVP (open bars) and AI (hatched bars) binding in the presence of control medium (–H7) was designated 100% and binding in the other experimental groups is expressed as percent of control (n = 3).
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Figure 8. Effect of a high extracellular glucose concentration on AVP and All induced 45Ca2+ efflux from VSMC. Confluent VSMC were incubated with either control medium (glucose 5 mM) or high glucose medium (glucose 20 mM) for 48 h and then preloaded with 45Ca2+. AVP (10−7) (open bars) and All (10−7 M) (hatched bars) stimulated 45Ca2+ efflux from VSMC was measured over a 5-min period and was expressed as percent total 45Ca2+ available at the time of stimulation. *P < 0.05 vs. all of the control media (n = 4).

mones rather than any change in the basal efflux rate. Furthermore, these data suggest that the magnitude of AVP and All receptor downregulation is paralleled by a proportionate decrease in intracellular signaling.

The process of Ca2+ mobilization in VSMC is closely linked to the intracellular mechanisms regulating cell contraction (44, 45). The next study therefore examined whether the glucose-induced downregulation of AVP and All receptors on VSMC was also associated with impaired VSMC contractile responses to AVP and All. Sparsely plated individual primary cultures of VSMC were incubated with one of the four test media for 48 h before examining VSMC contraction in response to AVP or All (10−6 M). Contraction was defined as a > 15% reduction in surface area, measured using microscopic digital planimetry. Spontaneous contraction of cells due to manipulation of the media was the same after preexposure to all test media. Moreover, median cell planar surface area was not significantly different after exposure to different test media, thus contractile responses were measured from a similar baseline surface area in the presence of all test media. In the presence of control medium, almost 50% of VSMC underwent a significant contractile response after a 15-min exposure to either AVP or All, compatible with data previously published from this laboratory (23, 24, 44). Results in the presence of the mannitol or L-glucose media were equivalent to control. However, after a 48-h exposure to a high D-glucose (20 mM) medium, the contractile response of VSMC to AVP and All was significantly impaired (Fig. 9). These data complement the signal-transduction studies in suggesting that the PKC-dependent downregulation of AVP and All receptors on VSMC exposed to a high glucose environment is associated with a significant depression of biochemical and physiological responses of VSMC to these pressor hormones.

Discussion

Several in vivo studies demonstrate that the early diabetic state is associated with a downregulation of diverse classes of pressor receptor on a variety of tissues (15–19). In spite of its potential pathophysiological significance with regard to vascular injury in diabetes mellitus, the factors responsible for receptor downregulation remain undefined. Hyperglycemia is the characteristic metabolic abnormality of diabetes mellitus and some studies demonstrate that an inverse relationship may exist between blood glucose concentration and hormone receptor density (16, 19). Treating rats with insulin to maintain euglycemia prevents glomerular All receptor downregulation (16). Furthermore, in diabetic patients, there is a significant negative correlation between platelet thromboxane A2 receptor number and glycosylated hemoglobin A1c levels (19), the latter being a measure of recent glycemic control. However, because of the multiplicity of factors that can influence the regulation of receptor expression, it has been impossible to examine the specific and independent effect of an elevated extracellular glucose concentration in vivo. To this purpose, the present study used an in vitro cell culture system to exclude other variables and uniquely demonstrates a direct and specific effect of high extracellular glucose concentrations to downregulate AVP and All receptors on vascular tissue. This effect is not attributable to changes in extracellular osmolality but is dependent on the metabolism of D-glucose. The receptor downregulation was apparent using glucose concentrations compatible with those attained in poorly controlled diabetic patients.

Analysis of receptor-binding kinetics reveals a similar effect of high extracellular glucose concentrations on both AVP and All binding to VSMC. The specific binding of both pressor hormones to VSMC was depressed due to a significant reduction in surface density of each receptor type rather than any significant change in receptor affinity. Of interest, similar characteristics for pressor receptor downregulation have been observed early in the course of experimental and human diabetes mellitus. In the rat glomerulus, All receptors are downregulated within 24 h of onset of diabetes mellitus (15, 16) and, in diabetic patients, the AVP (V1) and the thromboxane A2 receptor are downregulated on the surface of platelets (17, 19). In each instance, the diminished hormone binding resulted from decreased receptor density without a significant change in re-
Receptor affinity, an identical response in vitro to the effect of glucose on pressor receptors demonstrated in the present study. In addition, the magnitude of receptor downregulation due to diabetes mellitus in vivo (30–58%) is similar to the maximal glucose-induced response of ~40%. In each in vivo study, decreased pressor receptor number in diabetic animals and man could not be explained by previous receptor occupancy or homologous desensitization due to reciprocal changes in circulating levels of the relevant hormone (15–17, 19). Moreover, more detailed studies of the AII receptor revealed appropriate AII receptor regulation in diabetic rat glomeruli after pharmacological manipulation of the renin–angiotensin system (15, 16). These latter observations imply a direct effect of a metabolic consequence of the diabetic state on pressor receptor expression. One major consequence is hyperglycemia and the present in vitro study illustrates that a potent and specific effect increased glucose to downregulate AVP and AII receptors on cultured VSMC. This effect is both qualitatively and quantitatively similar to the effects of the diabetic state on these receptors in vivo. Although it is impossible to extrapolate directly from in vitro data to the in vivo situation, this striking correspondence does suggest that the previously documented effects of diabetes on pressor receptor kinetics in vivo may be a direct consequence of an elevated extracellular glucose concentration. The present results also provide information about the cellular mechanism whereby this effect of glucose may be mediated.

PKC may be important in regulating the expression of a diverse population of cell surface receptors in a variety of tissues (25–28). The present study demonstrates that high extracellular glucose concentrations induce a sustained increase in the activity of PKC in VSMC. This observation is compatible with those of others, which show a glucose-dependent activation of PKC in many tissues in vivo and in vitro (29–32). Using isolated glomeruli, Craven and coworkers recently demonstrated that elevated glucose concentrations increase the flux of glucose through an intracellular pathway that culminates in the enhanced de novo synthesis of DAG and concluded that this increase in DAG mass may contribute to glucose-induced activation of PKC (29, 32). Numerous studies in diabetic animals confirm that DAG levels are indeed elevated in a variety of tissues, including vascular tissue (29, 31, 32, 46). The present study lends support to the aforementioned hypothesis of Craven and coworkers by demonstrating that elevated extracellular glucose concentrations of the nonmetabolized glucose isomer, L-glucose, do not activate PKC, implying that a product of glucose metabolism, perhaps DAG, is required for glucose-induced activation of PKC in VSMC.

To explore the relevance of the sustained activation of PKC by glucose to the process of glucose-induced receptor downregulation in vascular tissue, glucose-induced PKC activation was prevented with H7, a relatively specific inhibitor of PKC activation (43). This maneuver almost completely prevented glucose-induced downregulation of the AVP or AII receptor, providing strong evidence that glucose-induced pressor receptor downregulation is dependent on the capacity for glucose to activate PKC in VSMC. It is important to note, however, that H7 is not totally specific for PKC and it therefore remains possible that the concomitant inhibition of other kinase systems could be contributing to the effects of H7 on glucose-induced receptor downregulation, as observed in the present study.

The present study demonstrates a slow onset and slow recovery of glucose-induced AVP and AII receptor downregulation, requiring hours. The recycling time for the AVP (V1) and AII receptors, however, is rapid in VSMC, ~15–20 min (47, 48). An effect of glucose to inhibit this process would have been more immediately apparent. Thus the slow downregulation of AVP and AII receptors by the high glucose environment and their slow recovery is more likely to reflect modulation of receptor biosynthesis. An effect of PKC on epidermal growth factor receptors has suggested this mechanism in the Swiss/3T3 line. It thus seems possible that the PKC dependency of the observed glucose-mediated receptor downregulation may inhibit the expression of receptor mRNA.

Diabetes-induced downregulation of pressor receptors may be relevant to the increased blood flow to many tissues that has been implicated as an important factor in the early pathogenesis of microvascular injury (8–10). The present study thus explored the biochemical and functional significance of glucose-induced AVP and AII receptor downregulation in VSMC. The results show that AVP- and AII-stimulated Ca2+ efflux from VSMC and the contractile response of individual VSMC to these agents are markedly attenuated after preexposing VSMC to a high glucose environment for 48 h. As in the effects on receptor density and PKC activation, these effects were specific for the high D-glucose medium. Moreover, the magnitude of glucose-induced AVP or AII receptor downregulation appeared to parallel the glucose-induced defect in AVP- and AII-stimulated intracellular signaling and contractility in VSMC. Although this does not prove causality, recent studies have demonstrated that there are few, if any, spare V1 or AII receptors on VSMC (47, 49), thus implying that the downregulation of either receptor on VSMC would be associated with a decrease in VSMC response. It should be emphasized that in addition to the role in receptor downregulation, glucose-induced PKC activation may also exert direct downregulatory influences on postreceptor signal transduction mechanisms in VSMC (23, 50–52). Recently, the glucose-induced PKC activation has been characterized in cultured VSMC (53).

In conclusion, the present study uniquely demonstrates that high extracellular glucose concentrations depress the expression of AVP and AII receptors on VSMC. This effect is dependent on glucose-induced PKC activation and is likely to be of physiological significance in view of the associated marked attenuation of VSMC functional responses to AVP and AII. This novel mechanism may contribute to pressor receptor downregulation in early diabetes mellitus, thereby directly implicating hyperglycemia in the pathophysiology of hemodynamically mediated vascular injury in this disease.

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References


